



INTERNATIONAL APPLICATION PUBLISHED UNDER THE PATENT COOPERATION TREATY (PCT)

(51) International Patent Classification ⁴ : A61K 45/02	A1	(11) International Publication Number: WO 88/03411 (43) International Publication Date: 19 May 1988 (19.05.88)
---	----	---

(21) International Application Number: PCT/US87/02998 (22) International Filing Date: 6 November 1987 (06.11.87) (31) Priority Application Number: 927,834 (32) Priority Date: 6 November 1986 (06.11.86) (33) Priority Country: US (71) Applicant: AMARILLO CELL CULTURE COMPANY, INC. [US/US]; 6666 Amarillo Boulevard West, Amarillo, TX 79106 (US). (72) Inventor: CUMMINS, Joseph, M. ; 3538 Barclay, Amarillo, TX 79109 (US). (74) Agent: LAMMERT, Steven, R.; Barnes & Thornburg, 11 South Meridian Street, 1313 Merchants Bank Building, Indianapolis, IN 46204 (US).	(81) Designated States: AT (European patent), AU, BB, BE (European patent), BG, BJ (OAPI patent), BR, CF (OAPI patent), CG (OAPI patent), CH (European patent), CM (OAPI patent), DE (European patent), DK, FI, FR (European patent), GA (OAPI patent), GB (European patent), HU, IT (European patent), JP, KP, KR, LK, LU (European patent), MC, MG, ML (OAPI patent), MR (OAPI patent), MW, NL (European patent), NO, RO, SD, SE (European patent), SN (OAPI patent), SU, TD (OAPI patent), TG (OAPI patent). Published <i>With international search report.</i> <i>With amended claims and statement.</i>
--	---

(54) Title: IMPROVED INTERFERON THERAPY

(57) Abstract

Neoplastic disease, hyperallergenicity, autoimmune disorders characterized by chronic tissue degenerative inflammation and immuno-resistant viral infections are treated by the administration of interferon at a dosage of about 0.1 to about 5 IU/lb per day by contacting said interferon with oral/pharyngeal mucosa. Interferon is administered in solution or in a novel solid unitary dosage form adapted to be dissolved in saliva when placed in the mouth.

FOR THE PURPOSES OF INFORMATION ONLY

Codes used to identify States party to the PCT on the front pages of pamphlets publishing international applications under the PCT.

AT	Austria	FR	France	ML	Mali
AU	Australia	GA	Gabon	MR	Mauritania
BB	Barbados	GB	United Kingdom	MW	Malawi
BE	Belgium	HU	Hungary	NL	Netherlands
BG	Bulgaria	IT	Italy	NO	Norway
BJ	Benin	JP	Japan	RO	Romania
BR	Brazil	KP	Democratic People's Republic of Korea	SD	Sudan
CF	Central African Republic	KR	Republic of Korea	SE	Sweden
CG	Congo	LI	Liechtenstein	SN	Senegal
CH	Switzerland	LK	Sri Lanka	SU	Soviet Union
CM	Cameroon	LU	Luxembourg	TD	Chad
DE	Germany, Federal Republic of	MC	Monaco	TG	Togo
DK	Denmark	MG	Madagascar	US	United States of America
FI	Finland				

IMPROVED INTERFERON THERAPY

BACKGROUND OF THE INVENTION

This invention relates generally to an improved method of treating diseases of immuno-pathologic etiology in warm-blooded vertebrates using interferon in low oral dosages. This invention also relates to the 5 use of interferon in low oral dosages to potentiate disease-corrective immune responses in warm-blooded vertebrates afflicted with immuno-resistant diseases characterized by apparent hyperactive or hypoactive immune system function.

10 "Interferon" is a term generically comprehending a group of vertebrate glycoproteins and proteins which are known to have various biological activities, such as antiviral, antiproliferative, and immunomodulatory activity at least in the species of 15 animal from which such substances are derived. The following definition for interferon has been accepted by an international committee assembled to devise a system for the orderly nomenclature of interferons: "To qualify as an interferon a factor must be a protein 20 which exerts virus nonspecific, antiviral activity at least in homologous cells through cellular metabolic processes involving synthesis of both RNA and protein." Journal of Interferon Research, 1, pp. vi (1980). "Interferon" as used herein in describing the present 25 invention shall be deemed to have that definition.

Since the first descriptions of interferon by Isaacs and Lindeman [See, Proc. Roy. Soc. London (Ser. B), Vol. 147, pp. 258 et seq. (1957) and U.S. Patent No. 3,699,222], interferon has been the subject of intensive 30

research on a worldwide basis. The literature is replete with publications concerning the synthesis of interferon, its proposed molecular characterizations, its clinical applications and proposed mechanisms of its antitumor, antiviral, and immune system activities.

Because of the intensity and disparate origins of research concerning interferon and its characteristics and uses, there exists a substantial lack of uniformity in such matters as classification of interferon types. There are also numerous, sometimes contradictory, theories concerning the mode of action of interferon in producing clinical effects.

Although originally isolated from cells of avian origin (chick allantoic cells), interferon production has been observed in cells of all classes of vertebrates, including mammals, amphibians, birds and reptiles. Interferon production by vertebrate cells is seldom spontaneous but is often readily "induced" by treatment of cells (in vivo or in vitro) with a variety of substances including viruses, nucleic acids (including those of viral origin as well as synthetic polynucleotides), lipopolysaccharides, and various antigens and mitogens.

Interferons have generally been named in terms of the species of animal cells producing the substance (e.g., human, murine, or bovine), the type of cell involved (e.g., leukocyte, lymphoblastoid, fibroblast) and, occasionally, the type of inducing material responsible for interferon production (e.g., virus, immune). Interferon has been loosely classified by some

researchers according to induction mode as either Type I or Type II, with the former classification comprehending viral and nucleic acid induced interferon and the latter class including the material produced as a lymphokine through induction by antigens and mitogens. More recently, the international committee devising an 5 orderly nomenclature system for interferon has classified interferon into types on the basis of antigenic specificities. In this newer classification, the designations alpha (α), beta (β), and gamma (γ) have been used to correspond to previous designations of 10 leukocyte, fibroblast, and type II (immune) interferons, respectively. Alpha and beta interferons are usually acid-stable and correspond to what have been called type I interferons; gamma interferons are usually acid-stable and correspond to what has been called type II 15 interferons. The international committee's nomenclature recommendations apply only to human and murine interferons. Journal of Interferon Research, 1 pp. vi (1980).

In its earliest applications, interferon was 20 employed exclusively as an antiviral agent and the most successful clinical therapeutic applications to date have been in the treatment of viral or virus-related disease states. It became apparent, however, that 25 exogenous interferon was sometimes capable of effecting regression or remission of various metastatic diseases. An overview of current clinical trials of interferon as an antiviral and antiproliferative therapeutic agent is contained in Interferon: In Vivo and Clinical Studies.

Volume 4, Eds: N. B. Finter and R. K. Oldham, Academic Press, New York, 1985.

The clinical agent of choice for the present is human leukocyte interferon, "mass-produced" by procedures involving collection and purification of vast quantities of human buffy coat leukocytes, induction with virus, and isolation from culture media.

In the work described above, interferon has been administered parenterally, i.e., intramuscularly and intradermally, with some successful topical and intranasal usages having been reported. It has seldom been administered intravenously because of substantial adverse effects attributable to "contaminants" in crude and even highly purified isolates.

As discussed above, there has been a significant research effort directed to the evaluation of therapeutic effects of interferon for a wide variety of diseases having an auto-immuno-pathologic basis. Before applicant's first report of successful oral administration of interferon in his U.S. Patent Application Serial No. 415,525 (now U.S. Patent 4,462,985), there was no recognition in the art of the potential offered by oral administration of interferon. The generally held belief was that interferon could not survive the digestive conditions of the upper alimentary canal.

Since applicant's first disclosure of the immunotherapeutic benefit achievable via oral administration of interferon of heterologous mammalian species, he has continued to investigate the efficacy of

orally administered interferon. In U.S. Patent No. 4,497,795, issued February 5, 1985, applicant described and claimed the use of interferon administered orally or via intravenous administration to stimulate appetite and feed efficiency of bovine and porcine species. More recently applicant has described in now pending U.S. 5 applications the use of interferon at dosages less than about 5 IU per pound of body weight for increasing feed efficiency and food utilization in warm-blooded vertebrates, for preventing and treating shipping fever, and for enhancing vaccine efficiency.

10 Human alpha-interferon has been marketed under the trademark Agriferon® by Immunomodulator Laboratories, Inc. ("IML") of Stafford, Texas for veterinary use in Texas since February 1985. The product is sold for oral administration to cattle to 15 promote growth and feed efficiency and to prevent or treat viral respiratory infections. IML began selling an alpha-interferon product for horses in 1986. Both products are sold under a license of my U.S. Patent 4,462,985.

20

SUMMARY OF THE INVENTION

25 Interferon contacting the oral and/or pharyngeal mucosa, in amounts of less than 5 IU/lb of body weight per day is consistently effective to potentiate disease-corrective immune responses in vertebrates afflicted with immuno-resistant disease states characterized by apparent hyperactive or

30

hypoactive immune system function. Treatment in accordance with the present invention has been shown to effect remission of neoplastic disease, hyperallergenicity, immuno-resistant or immuno-debilitating viral infections and autoimmune disorders characterized by chronic tissue degenerative inflammation.

DETAILED DESCRIPTION OF THE INVENTION

The clinical agent of choice for use in the present invention is human leukocyte interferon (human alpha-interferon), "mass-produced" by procedures involving collection and purification of quantities of human buffy coat leukocytes, induction of interferon production with virus, and isolation of culture media. (See "Preparation of Human Alpha-Interferon" below.) Also acceptable for use in accordance with present intention are human alpha-interferon products produced by recombinant DNA technology and now commercially available from Schering-Plough (as Intron®) and Hoffmann-LaRoche (as Roferon®) and approved by the FDA for treatment (parenterally) of hairy cell leukemia of man. Such recombinant interferon products are believed to be particularly effective when used in combination. Gamma interferon is also available by recombinant technology and is presently undergoing clinical trials by Genentech and others. Fibroblast interferon (beta-interferon) can be prepared in accordance with Example 1 in applicant's U.S. Patent No. 4,462,985

issued July 31, 1984, the disclosure of which is hereby expressly incorporated by reference.

Interferon of human and murine origins has been quantified in the art in terms of International Units ("IU"). As used herein, a "unit" of interferon (to be distinguished from "IU") shall mean the reciprocal of a dilution of interferon-containing material that, as determined by assay, inhibits one-half the number of plaques of a challenge virus, the challenge virus being the vesicular stomatitis virus ("VSV"). So quantified a "unit" of interferon is routinely found to be about one-tenth the quantity of interferon represented by one "IU." In other words, for the purpose of defining the present invention, 1 unit \approx 0.1 IU.

The present invention relates to an improved method of treatment of immuno-resistant disease states with interferon. The present invention is directed to the treatment of diseases in warm-blooded vertebrates, particularly certain diseases which the immune system of many species is poorly equipped to handle, as evidenced by either a lack of disease defeating response and/or an apparently misdirected immune response resulting in a chronic tissue degenerative inflammatory condition or other physical complications. While there has been a significant research effort directed to the use of interferon for treatment of such diseases, reported results, although positive overall, have been inconsistent. The principle reason for such inconsistency in view of my most recent research efforts is that earlier investigators have failed to define optimum dosage and route of interferon administration.

The present invention is based on applicant's discovery that interferon can be used as a consistently effective therapeutic agent for treatment of diseases having an immunopathologic basis - characterized by inadequate immune response and persistence of the disease or by an apparent hyperactive immune response resulting in tissue degenerative inflammatory conditions and related physical manifestations. Applicant has found that interferon, contacting the oral and pharyngeal mucosa in amounts from about 0.01 to about 5 IU/lb of body weight per day, is consistently efficacious for the treatment of diseases to which the immune system of many warm-blooded vertebrates does not effectively respond.

Disease conditions treated in accordance with the present invention include apparent autoimmune disorders characterized by a chronic tissue degenerative inflammatory condition. Diseases so characterized include multiple sclerosis, rheumatoid arthritis, stomatitis, and lupus erythematosus. Treatment of such disease is in accordance with the present invention comprises administering interferon at a dosage of 0.01 to about 5 IU/lb per day in a dosage form adapted to promote contact of said dosage of interferon with the oral and pharyngeal mucosa of said animal. Preferably, the dosage of interferon is from 0.1 to about 4.0 IU/lb per day, more preferably 0.5 to about 1.5 IU/lb of body weight per day. Alpha interferon, derived from tissue culture or by recombinant DNA techniques, is a preferred therapeutic agent in accordance with this invention.

Alpha interferon can be administered alone or in combination with beta interferon or gamma interferon.

It is critical that the interferon be administered in a dosage form adapted to assure maximum contact of the interferon in said dosage form with the oral and pharyngeal mucosa of the human or animal
5 undergoing treatment. Contact of interferon with the mucosa can be enhanced by maximizing residence time of the treatment solution in the oral or pharyngeal cavity. Thus, best results seem to be achieved in human patients when the patient is requested to hold said
10 solution of interferon in the mouth for a period of time. Contact of interferon with the oral and pharyngeal mucosa and thereafter with the lymphatic system of the treated human or animal is unquestionably
15 the most efficient method administering immunotherapeutic amounts of interferon.

Another disease condition responding to treatment in accordance with the present invention is neoplastic disease. Thus, the administration of interferon in accordance with the above description can, alone or in combination with other drugs or therapy,
20 help effect remission of cancers such as malignant lymphoma, melanoma, mesothelioma, Burkitt lymphoma and nasopharyngeal carcinoma and other neoplastic diseases, especially those of known or suspected viral etiology.
25 Based on the results observed to date, it is believed that applicant's presently described method of treatment will similarly help effect remission of Hodgkin's Disease and leukemia.

Other disease conditions responding to treatment in accordance with the present invention are infectious diseases of viral origin in, for example, human, avian, porcine, canine and feline species. Significantly, viral infection typically exhibiting persistent resistance to treatment have shown a dramatic response to treatment with interferon in low doses contacting the oral and pharyngeal mucosa of infected patients. Beneficial results have been attained utilizing the present method to treat dogs having canine parvovirus and canine herpesvirus infections. Further, feline leukemia and feline infectious peritonitis have been shown to be particularly susceptible to treatment with alpha interferon and beta interferon in accordance with this invention.

Exemplary of human viral infections showing remarkable response to treatment in accordance with the present invention are infections of human rhinovirus (common cold), herpes simplex I virus (cold sores) and human papovavirus (warts). Based on treatment results to date, it is expected that contact of interferon at low dosage with the oral and pharyngeal mucosa will provide an effective treatment for Acquired Immune Deficiency Syndrome (AIDS) and disease conditions having the herpes simplex II virus as the causative agent. A patient experiencing a condition of viral myocarditis has responded favorably to the present treatment. Warts often dissipate within six to eight weeks after initiating treatment in accordance with this invention.

Other afflictions responding to contact of low dosage interferon are hyperallergenic conditions such as asthma. One "side effect" noted by patients treated in accordance with this invention is improved skin complexion. Thus, administration of interferon in dosages of about 0.01 to about 5 IU/lb of body weight per day is effective to treat acne, specifically and improve human skin complexion generally.

Further, stimulating the immune system by oral contact with low dosage interferon is believed to assist the body in fighting bacterial infection. Treatment in accordance with this invention alone or in combination with therapeutic amounts of antibiotics can be especially effective in knocking down infections of antibiotic resistant microorganisms.

Administration of interferon in accordance with the present invention is preferably continued until the symptoms of the disease condition being treated subside. This can range from a period of one day, for example, where a human rhinovirus is the disease causative agent, to a period of up to six months for treatment of neoplastic disease. Rheumatoid arthritis patients are pain free within 2 to 10 days of initiating treatment in accordance with the present invention. However, treatment of that disease is preferably conducted by administration of interferon for up to about three (3) months.

Daily dosage of interferon can be administered as a single dosage or, preferably, it is divided and administered in a multiple-dose daily regimen. A

staggered regimen, for example one to three days treatment per week or month, can be used as an alternative to continuous daily treatment.

Interferon can be administered in accordance with this invention in either a liquid (solution) or solid dosage form. Thus interferon can be administered
5 dissolved in a buffered aqueous solution typically containing a stabilizing amount (1-5% by weight) of blood serums. Exemplary of a buffered solution suitable as a carrier of interferon administered in accordance
10 with this invention is phosphate buffered saline prepared as follows:

A concentrated (20x) solution of phosphate buffered saline (PBS) was prepared by dissolving the following reagents in sufficient water to make 1,000 ml of solution: sodium chloride, 160 grams; potassium chloride, 4.0 grams; sodium hydrogen phosphate, 23 grams; potassium dihydrogen phosphate, 4.0 grams; and optionally phenol red powder, 0.4 grams. The solution is sterilized by autoclaving at 15 pounds pressure for 15 minutes and then diluted with additional water to a single strength concentration prior to use.
15
20

Alternatively the interferon can be formulated into flavored or unflavored solutions or syrups using a buffered aqueous solution of interferon as a base with added caloric or non-caloric sweeteners, flavor oils and pharmaceutically acceptable surfactant/dispersants.
25

It is also contemplated by the present invention to provide interferon in a solid dosage form such as a lozenges adapted to be dissolved upon contact

with saliva in the mouth with or without the assistance of chewing. Such a unitary dosage form is formulated to release about 1 to about 1500 IU of interferon upon dissolution in the mouth for contact with the oral and pharyngeal mucosa. Thus a unitary dosage form of interferon in accordance with this invention can be prepared by art-recognized techniques for forming compressed tablets such as chewable vitamins.

5 Similarly, interferon can be incorporated into starch-based gel formulations to form a lozenge which will dissolve and release interferon for contact with the oral mucosa when held in the mouth. Solid unitary dosage forms of interferon for use in accordance with the present invention can be prepared utilizing art recognized dosage formulation techniques. The pH of such formulations can range from about 4 to about 8.5.

10 15 Of course, in processing to such unitary dosage forms one should avoid heating a pre-dosage form formulation, after addition of interferon, above about 50° Centigrade.

Preparation of Human Alpha-Interferon

20

Human alpha-interferon can be prepared through the following procedure, commonly referred to as the Cantell procedure. The process begins with packs of human leukocytes, obtained in this case from the Gulf Coast Regional Blood Center, Houston, Texas. The buffy coats in these packs are pooled into centrifuge bottles, and then are diluted with 0.83% ammonium chloride. The mixture is incubated for 15 minutes with intermittent

25 30

shaking, and is then centrifuged for 20 minutes at 2000 rpm. The supernatant is discarded, and the cell pellets are resuspended with a minimal volume of sterile phosphate buffered saline (PBS). The mixture is then diluted with ammonium chloride and centrifuged. The supernatant is again discarded, and the remaining cell 5 pellets are resuspended with a minimal volume of a tissue culture medium such as Minimal Essential Medium (MEM), available from KC Biological. The cell concentration is determined with a Coulter counter.

10 Interferon induction takes place in glass or plastic bottles. The induction medium contains MEM, 75mM Hepes (available from Calbiochem), 75mM Tricine (available from Sigma Chemical Co.), human agamma serum (18mg/ml), and gentamycin sulfate (from M.A. Bioproducts; 50mcg/ml). The cells are added to the 15 induction vessels at a final concentration of about 5 to 10 million cells per milliliter. The induction vessel is incubated in a 37°C water bath, and interferon alpha is added as a primer.

20 After two hours, Sendai virus is added to the induction mixture. This causes alpha interferon to be produced in the supernatant by the leukocytes. After a 12-18 hour incubation time, the induction mixture is centrifuged. The cells are discarded, and the supernatant is then purified.

25 The crude interferon is chilled to 10°C or below in an ice bath. Five molar potassium thiocyanate is added to obtain a final concentration of 0.5M. This solution is stirred for 15 minutes, and then its pH is

lowered to 3.3 by adding hydrochloric acid. The mixture is then centrifuged at 2800 rpm for 30 minutes, and the supernatant is discarded.

The pellets are then resuspended in 95% ethanol and are stirred for 15 minutes. This suspension is centrifuged at 2800 rpm for 20 minutes, and the pellets are discarded. The pH of the supernatant is then adjusted to 5.8 with sodium hydroxide. The mixture is stirred for 10 minutes, and then centrifuged at 2800 rpm for 20 minutes. The pellets are discarded. The pH of the supernatant is then adjusted to 8 with sodium hydroxide. This solution is stirred for 10 minutes, followed by centrifugation at 2800 rpm for 20 minutes. The supernatant is discarded, and the pellets are resuspended with 0.5M potassium thiocyanate in a 0.1M sodium phosphate buffer. This suspension is stirred at 15 4°C.

Next, the suspension is centrifuged at 2800 rpm for 20 minutes, and the pellets are discarded. The pH of the supernatant is adjusted to 5.3 with hydrochloric acid. After stirring for 10 minutes and centrifugation, the pH of the supernatant is adjusted to 2.8 with hydrochloric acid, followed by further stirring for 20 minutes. This mixture is centrifuged at 2800 rpm, and the resulting pellet is purified human alpha-interferon.

The pellet is resuspended with 0.5M potassium thiocyanate in 0.1M sodium phosphate buffer, having a pH of 8.0. It is then dialyzed against PBS at 4°C, with two changes of PBS. This mixture is then centrifuged and the precipitate is discarded. The remaining

purified alpha interferon is sterilized by filtration through a 0.2 micron filter. A human alpha-interferon is produced in accordance with this procedure by Immuno Modulators Laboratories, Inc., Stafford, Texas, and sold under the trademark Agriferon® for use in cattle and Equiferon® for use in horses.

5 Other procedures known to those skilled in the art are available for making interferons, such as human alpha-interferon and human gamma-interferon. For example, U.S. Patents 4,376,821 and 4,460,685 disclose methods of making human gamma-interferon. A method of
10 making bovine fibroblast (beta) interferon is disclosed in applicant's U.S. patent 4,462,985.

Clinical Studies

15 Tables 1-4 below summarize the results of clinical studies of the administration of interferon by veterinarians orally to 137 dogs and cats as of November, 1985. The studies were conducted with both human alpha-interferon and bovine beta-interferon.

20 Tables 1-4 compare survival rates of pets with feline leukemia virus-associated diseases or canine parvovirus disease. Unequal numbers of pets were treated with each type of interferon; bovine beta-interferon was given to 78 pets and human alpha-interferon was given to 59 pets.

25 Bovine beta-interferon was produced in flasks of confluent monolayers of bovine fetal kidney (BFK) cells. Culture supernatant was harvested 24 hours after bluetongue virus induction of BFK cells. The

supernatant was dialyzed 24 hours in a pH 2.0 buffer and for another 24 hours in a PBS (pH 7.4) buffer before interferon assay. Procedures for the assay and characterization of bovine beta-interferon were essentially as described by Rosenquist and Loan, American Journal of Veterinary Research, 28; 619-628, 5 1967. Interferon titers as "units" were expressed as the reciprocals of the dilutions that provided a 50% reduction in the number of VSV plaques as compared with the number in control cultures. The BFK cell culture interferon produced by this method had an average titer 10 of 7,000 units per milliliter. Dogs were given bovine beta-interferon, 5-10 ml/dose, at least three times/day after a diagnosis of CPV disease. Cats positive by ELISA for feline leukemia virus and exhibiting clinical signs of disease were given 1 ml/10 lb of body weight 15 2-3 times daily for five days. After a five-day interval, cats were retreated at least once for another five days.

Human alpha interferon was obtained from IML, Inc. of Houston, Texas. Cases were treated with lot 20 AO26 applied at 6×10^6 IU/ml. Lot AO26 of human alpha interferon was diluted 1:150 in Eagles' minimum essential medium (MEM) and used as the stock solution from which 1 ml was further diluted 1:1000 with 1 liter of MEM for treatment. The usual dose of human alpha 25 interferon was 4 IU/lb body weight given at least three times daily after a diagnosis of CPV disease was made. For feline leukemia, cats were treated with human alpha-interferon 2-3 times daily for five days as reported for bovine beta-interferon.

Significantly ($P<0.05$) more cats lived six and twelve months after diagnosis and treatment for feline leukemia virus if alpha-interferon was given, compared to treatment with bovine beta-interferon. Significantly ($P<0.05$) more dogs survived CPV disease when given alpha interferon (92%) compared to those dogs given bovine beta-interferon (69%).

5

TABLE 1

10

Summary of Survival Date from clinically ill cats positive for FeLV.

<u>Treatment</u>	<u>Months After Treatment</u>		
	<u>1</u>	<u>6</u>	<u>12</u>
Human alpha-IFN	25/33	21/32*	19/31*
Bovine IFN	26/36	15/36	13/36

15

Numerator = no. alive; denominator = no. treated.

*Cats given human alpha-IFN had significantly ($P<.05$) higher survival rates at 6 and 12 months after treatment than cats given bovine IFN. Significance was determined by Chi-Square test.

25

30

TABLE 2

Percent survival of clinically ill cats positive for FeLV.

Treatment	Months After Treatment		
	1	6	12
Human alpha-IFN	76%	66%*	61%*
Bovine IFN	72%	42%	36%
<u>Historical Control <50%</u>		<30%	--

Numerator = no. alive; denominator = no. treated.

*Cats given human alpha-IFN had significantly ($P<.05$) higher survival rates at 6 and 12 months after treatment than cats given bovine IFN. Significance was determined by Chi Square test..

15

TABLE 3

Response of CPV disease to treatment with bovine interferon or human alpha-interferon, by veterinarian.

	Attending Veterinarian	Bovine IFN Beta		Human Alpha IFN	
		Lived	Died	Lived	Died
S	16/21	5/21	14/16	2/16	
M	6/11	5/11	7/7	0/7	
R	7/10	3/10	3/3	0/3	
25	29/42(69%)	13/42(31%)	24/26(92%)	2/26(8%)	

Dogs treated with human alpha-interferon had a significantly ($P<.05$) higher survival rate compared to dogs treated with bovine IFN. Significance between groups was determined by Chi Square test.

30

-20-

TABLE 4

Treatment days for CPV disease.

	<u>No. of Treatment</u>	<u>Average Days</u>	<u>No. Treatment</u>	<u>Survival Rate</u>
	<u>Treatment</u>	<u>Days</u>	<u>Days*</u>	<u>SD**</u>
5	Bovine IFN	42	3.31	1.95
	Human alpha IFN	26	2.75	0.92

*Calculated on surviving dogs only.

**Standard deviation of the mean treatment days.

10.

Canine Herpesvirus Challenge of Newborn Dogs

15 Canine herpesvirus infection of dogs less than one week of age are invariably fatal, but older pups usually survive. Interferon has been successfully used to treat viral infections of many species. These studies were conducted to assess the efficacy of interferon in canine herpesvirus (CHV) inoculated puppies.

20 Five (5) pregnant bitches were obtained from a USDA licensed supplier and were housed in a USDA approved research facility in Canyon, Texas. After the pups whelped, they were inoculated with 6.3 log 10 units of virulent CHV obtained from Dr. Richard Mock of the 25 Texas A&M University Veterinary Medical Diagnostic Laboratory (TVMMDL) in Amarillo, Texas. Human alpha-interferon (HAI) or placebo was given to pups orally as treatment in an effort to increase the

30

survival rate of the CHV inoculated pups. Each litter was divided into control and treated animals. The procedures and schedule for each litter are discussed below. All dead animals were necropsied at TVMDL, Amarillo, Texas.

5 LITTER 1:

Nine (9) pups were inoculated orally with CHV on the day of birth. Interferon was given at 6-10 units (1X), or ten times the dosage at 60-100 units (10X). Three pups were given 0.5 ml placebo, 3 pups were given 10
10 0.5 ml HAI (1X), and 3 pups were given 0.5 ml of a 10X concentrate of HAI orally twice daily for 7 days (if they lived that long). The 3 controls died 5, 6, and 8 days after CHV inoculation. The 3 pups given HAI (1X)
15 lived 7, 7, and 9 days and the 3 pups given 10X HAI lived 6, 7, and 7 days after CHV inoculation.

Pups given HAI (1X) lived an average of 1.3 days longer than controls, but the longer survival time was not statistically significant. The higher dosage,
20 HAI (10X), did not provide a survival benefit over the lower dosage, but instead pups given the higher dosage died, on the average, one day sooner.

LITTER 2:

25 Eight (8) pups were inoculated with CHV orally 2 days after birth. Interferon was given at 6-10 units (1X), or ten times the dosage (10X), or 1/10th the dosage (1/10X). All interferon was given orally after dilution in PBS. Two (2) pups were given 0.5 ml PBS, 2
30

were given 0.5 ml HAI at 1/10th concentration, 2 were given HAI (1X), and 2 were given a 10X concentrate of HAI. All treatments were given orally twice daily for 5 days starting 1 day before CHV inoculation. The 2 controls died <1 and 9 days after CHV inoculation and the HAI treated (1/10th dose, full dose, 10X dose) pups died 8 and 9, 5 and 9, 8 and 8 days after CHV inoculation, respectively.

No benefit from treatment at any dosage was seen. The death of a control pup within a day after CHV inoculation was probably not related to CHV inoculation.

10

LITTER 3:

Nine (9) pups were inoculated with CHV orally 3 days after birth. Two pups were given 0.5 ml PBS, 2 were given 0.5 ml HAI (1X), 2 were given 1/10th dose HAI, and 3 were given 2 IU of recombinant human alpha-interferon from Schering-Plough. All treatments were given orally twice daily for 5 days starting two days before CHV inoculation. Both pups given HAI (1X) survived until necropsied 19 days after CHV inoculation. One control pup died 14 days after CHV inoculation and 1 survived until necropsied 19 days after CHV inoculation. Pups given 1/10th dosage of HAI died 8 and 13 days after CHV inoculation. Only one pup given recombinant human alpha-interferon died (12 days after CHV); the other 2 pups survived until necropsied 19 days after CHV inoculation.

25

20

30

These pups, inoculated 3 days after birth, did not develop an overwhelming CHV infection (only 1 of 2 controls died). A 1/10th dose of HAI did not protect either pup but both HAI (1X) treated pups survived.

LITTER 4:

Fourteen (14) pups were inoculated orally with CHV 2 days after birth. Seven (7) pups were given PBS and 7 pups were given HAI (1X) orally twice daily for 7 days (if they lived that long) starting 2 days after CHV inoculation. The 7 controls died 1, 5, 7, 8, 8, 9, and 9 days after CHV inoculation. One of the HAI (1X) treated pups survived and the other 6 pups died 1, 6, 8, 9, 9, and 12 days after CHV inoculation. The deaths of 2 pups only 1 day after CHV inoculation were probably not related to CHV inoculation.

One HAI (1X) treated pup lived 3 days beyond the last surviving control and one HAI (1X) treated pup lived 2 weeks (until necropsied) beyond any treated control pup. Average survival time of interferon treated pups was longer than control survival time, but not significantly so.

LITTER 5:

Six (6) pups were inoculated orally with CHV 2 days after birth. Three (3) pups were given PBS and 3 were given HAI (1X) orally once daily starting 5 days after CHV inoculation. The 3 controls died 6, 6, and 7 days after CHV inoculation. One of the HAI (1X) treated pups survived (until necropsied) and the others died 8 and 9 days after CHV inoculation.

All interferon treated pups lived longer than any of the control pups. Treatment with HAI (1X) did not begin until 5 days after CHV inoculation, yet survival was significantly ($P < .05$) prolonged.

In summary, on the average, puppies treated with human alpha-interferon had longer survival times

-24-

and enhanced survival rates compared to littermate controls, after canine herpesvirus challenge. A total of 7 puppies (1 control and 6 interferon treated) survived the normally fatal CHV inoculation. The data is summarized in the Table 5 below.

5

TABLE 5
Summary of Canine Herpesvirus Data

	<u>Litter</u>	<u>No. of Pups</u>	<u>Dosage</u>	<u>Average * Survival Time (Days)</u>	<u>Survivors</u>
10	1	3	control	6.33	0
	1	3	HAI 1X	7.67	0
	1	3	HAI 10X	6.67	0
	2	2	control	4.5**	0
	2	2	HAI 1/10th	8.5	0
	2	2	HAI 1X	7.0	0
15	2	2	HAI 10X	8.0	0
	3	2	control	14.0	1
	3	2	HAI 1/10th	10.5	0
	3	2	HAI 1X	-	2
20	3	3	recombinant IFN	12.0	2
	4	7	control	6.7	0
	4	7	HAI 1X	7.5	1
	5	3	control	6.3	0
	5	3	HAI 1X	8.5	1

25 * dead dogs survival time; living puppies not calculated.

** includes one pup living only one day beyond CHV inoculation.

Treatment Of Nasal Solar Dermatitis

Three cases of nasal solar dermatitis (collie nose) cleared after human alpha-interferon treatment of 1 unit/lb body weight orally and topical treatment (a few ml at 20 units/ml).

5

Treatment of Canine Lupus Erythematosus

Two cases diagnosed as canine lupus erythematosus were cured by human alpha-interferon treatment. A 2 year-old Lhasa apso male had been treated with prednisolone for 1 year for 3 dermatological lesions on the abdomen and prepuce. The flat glistening lesions were continually licked by the dog. Within 1 week of oral human alpha-interferon treatment (1 unit/lb body weight daily for 5 days, then after 1 week, treatment was repeated for 5 days) 2 lesions completely healed and the third lesion was reduced to 1/2 its original size. Within 4 weeks, the lesions were all completely healed and all therapy ceased. One year later, a skin lesion reappeared but promptly healed after interferon treatment was repeated. The skin lesions have not reappeared in the past 10 months.

A 6 year-old spayed female chihuahua cross had a spider shaped (4 cm by 2 cm approximately) skin lesion on the abdomen. The lesion was flat, glistening and pruritic. Six weeks of prednisolone treatment resulted in complete healing. The following summer, the lesion reappeared and was treated with oral human

30

alpha-interferon at about 1 unit/lb body weight daily for 5 days. Within 5 days the lesion was reduced to 1/3 its original size and completely disappeared within 10 days. The lesion has not reappeared in the past year.

Treatment Of Feline Infectious Peritonitis

5

Table 6 shows the results of 17 cases of feline infections peritonitis (FIP) as diagnosed by practicing veterinarians. Human alpha-interferon (IFN) treatment resulted in a significantly greater survival rate than treatment with bovine beta-IFN.

10

10

TABLE 6

Survival of clinically ill cats diagnosed as FIP

15

20

Treatment	No. Cats Treated	Alive	Dead	Survival Rate
Human alpha-IFN	11	10	1	91%
Bovine beta-IFN	6	3	3	50%
Total	17	13	4	76%

Cats given human alpha-IFN had a significantly ($P=.0574$) greater survival rate than cats given bovine beta-IFN.

25

30

Human Treatment With Exogenous
Human Alpha-Interferon

Human patients were treated with human alpha-interferon in the therapy of acute rheumatoid arthritis, multiple sclerosis, asthma, acne, malignant lymphoma, mesothelioma, and aphous stomatitis. Therapy consisted of oral administration of 0.7 IU per lb. of patient body weight twice daily, once in the morning and once in the evening. None of the patients noted any fever or anorexia associated with the administration of alpha interferon. Interferon was administered in a buffered solution having a concentration such that a single dosage could be administered in a volume of about 1 to about 20 ml of liquid. Each patient generally retained the interferon solution in his mouth for a period of time up to about one minute. After that time the solution was either swallowed or discharged from the patient's mouth.

Two patients suffering from rheumatoid arthritis were treated -- a Caucasian male age 44 and a Caucasian female age 44. The male patient was pain free in 7 days, and the female was pain free in 10 days. They were both continued on the oral interferon for 21 days total and have remained asymptomatic.

It has been found that recurrence of a treated arthritic condition can be minimized if treatment in accordance with the present invention is continued over a period of up to about three months.

A 30-year-old Caucasian female nurse afflicted with multiple sclerosis and who had had an extensive

neurologic workup at City of Hope Hospital in Los Angeles received treatment in accordance with the present invention for 21 days. The patient has had no recurrence of her neurologic symptoms for the past nine months.

5 A 42-year-old Caucasian male diagnosed to have a malignant lymphoma had completed chemotherapy with dismal results and was considered terminal. He was treated for three weeks with oral interferon. Six months after starting treatment he was released by his oncologist as free of the disease.

10 An 82-year-old Caucasian female was diagnosed to have mesothelioma. Presently there is no effective treatment for that disease and only a 9-month average survival rate is predicted. During her treatment with human alpha-interferon she had thoracentesis on two 15 occasions for plural effusion. Otherwise, the patient has been active and has survived for 43 months.

20 A 32-year-old Asian male with aphous stomatitis was treated for two weeks with human alpha-interferon in accordance with the present invention. There has been no recurrence of the ulcers over the last six months since treatment was completed.

25 BKC is a 29 year-old Caucasian female and KKJ is a 20 year-old Caucasian female. Both are afflicted by acne-like skin blemishes at the time of their monthly menstrual cycle. Oral human alpha-interferon given at about 1 unit/lb of body weight for 3 days prior to the time of their cycle reduces the severity and number of skin blemishes.

Treatment Of Warts In Humans
With Bovine Alpha-Interferon

MAH, a 38 year-old Caucasian female,
had 7 warts on the middle finger of her right hand.
After 9 months duration, medical treatment was sought,
5 and liquid nitrogen was applied by a dermatologist.
Only one wart on the finger regressed after treatment.
Three warts coalesced to create a large wart area that,
over the next year, acquired a roughly 12 millimeter
square shape. Oral bovine alpha interferon treatment
10 was started at a dosage of 6 ml daily for 6 consecutive
days. The concentration of alpha-interferon was 30
units/ml; it was derived from the nasal secretions of
cattle infected with infectious bovine rhinotracheitis
virus. All the warts completely regressed within 6
15 weeks of the first dose of interferon.

Interferon Dosage Formulations

(1) Lozenge

20 A starch gel-based lozenge containing
interferon is prepared by combining 150 grams of
sucrose, 550 ml phosphate buffered saline, and 250 grams
of a cold-water-soluble starch such as that described in
U.S. Patent 4,465,702, heating that mixture with
25 stirring to a temperature of about 75°C, cooling the
mixture to about 30°C and thereafter blending into the
paste-like mass 50 ml of phosphate buffered saline PBS
containing human alpha interferon at a concentration of

250 IU/ml. The mixture is then formed into multiple portions of about 5 to about 10 grams each which set upon standing under drying conditions to a starch candy gel-like consistency. The lozenges thereby produced can be administered to a patient singly or in combination. The patient is instructed to hold the lozenge in his 5 mouth until it is completely dissolved to release the interferon component for contact with the oral mucosa.

(2) Chewable Vitamin

A chewable vitamin formulation is prepared, for 10 example, according to the description of U.S. Patent 3,857,939 by coating one or more components thereof prior to tabletting with an interferon solution in an amount sufficient to provide about 1 to about 1500 units of interferon in each chewable vitamin tablet.

15

(3) Mouthwash

A mouthwash formulation is prepared in accordance with the present invention by combining 850 ml PBS, 100 ml of glycerin, 50 grams of dextrose, and a 20 mixture of 0.3 ml of a flavor oil pre-mixed with 30 ml of a palatable, pharmaceutically acceptable surfactant/dispersant having an HLB from about 15 to about 25 and 50 ml of a PBS solution of interferon (concentration 120 IU/ml). The formulation contains 25 interferon at a concentration of about 120 IU per 20 ml dosage. The patient is asked to hold a 20 ml volume of the mouthwash in his mouth, optionally gargling with the same, for a period of about 15 seconds to about one minute.

30

(4) Syrup

Interferon is added to a commercial cough syrup formulation in an amount sufficient to provide an interferon containing syrup formulation having about 1 to about 1500 IU of human interferon per tablespoon of syrup.

5

(5) Effervescent Tablet

A tableting mixture comprising a pharmaceutically acceptable alkali metal carbonate or bicarbonate, an organic acid such as citric acid, human interferon (preferably dispersed on a suitable organic or inorganic carrier therefor) in an amount sufficient to provide a per tablet dose of about 1 to about 1500 units of interferon per dose, and further including suitable tableting excipients such as lubricants and binders, is compressed into a unitary dosage form of interferon. The compressed tablet effervesces upon contact with water to release interferon to the resulting buffered solution. The dosage of interferon is readily available in solution for contact with the oral pharyngeal mucosa of a patient in need of said dosage of interferon.

25

30

CLAIMS

1. An improved method for administering interferon to a warm-blooded vertebrate for immunomodulation of said vertebrate, said method consisting essentially of contacting interferon with the oral and pharyngeal mucosa of said vertebrate at a dosage of about .01 to about 5 IU/lb. (about .022 to about 11 IU/kg).

2. The method of claim 1 wherein the administered interferon is human alpha-interferon or human beta-interferon.

3. The method of claim 2 wherein the interferon is administered at a dosage of about 0.1 to about 4.0 IU/lb (about 0.22 to about 8.8 IU/kg) per day.

4. The method of claim 1 wherein the interferon is administered to treat a disease selected from the group consisting of autoimmune disorder, neoplastic disease, viral infection, bacterial infection and hyperallergenicity.

5. The method of claim 4 wherein human alpha-interferon is administered to a human patient at a dosage of about 0.5 to about 1.5 IU/lb daily until the patient becomes asymptomatic of the disease.

6. The improvement of claim 1 wherein the administered interferon is human beta-interferon.

7. The improvement of claim 4 wherein human interferon is administered to a human patient as an aqueous solution in a dosage of about 0.5 to about 1.5 IU/lb per day and wherein said solution of interferon is

held in the patient's mouth for a period of time sufficient to allow contact of the interferon by the oral mucosa of said patient.

8. The method of claim 4 wherein the treated disease is an autoimmune disorder selected from the group consisting of multiple sclerosis, rheumatoid arthritis, nasal solar dermatitis, stomatitis and lupus erythematosus

9. The method of claim 4 wherein the neoplastic disease is selected from the group consisting of malignant lymphoma, melanoma, mesothelioma, Burkitt lymphoma and nasopharyngeal carcinoma.

10. The method of claim 9 wherein the neoplastic disease is selected from the group consisting of Hodgkin's disease and leukemia.

11. The method of claim 9 wherein the dosage of interferon is divided and administered in a multiple-dose daily regimen.

12. The method of claim 9 wherein neoplastic disease in humans is treated by administering human beta-interferon.

13. An improved method of treatment of infectious disease of viral origin in human, canine and feline species consisting essentially of administering interferon at a dosage of about 0.01 to about 5 IU/lb per day in a dosage form adapted to promote contact of said dosage of interferon with the oral and pharyngeal mucosa of said species.

14. The method of claim 13 wherein the dosage is about 0.1 to 4.0 IU/lb per day.

15. The method of claim 13 wherein the interferon is selected from alpha-interferon and beta-interferon.

16. The method of claim 13 wherein the interferon is human alpha-interferon.

17. The method of claim 13 wherein human alpha-interferon is administered at dosage of about 0.5 to 1.5 IU/lb of body weight per day.

18. The method of claim 13 wherein a human is treated for a viral infection selected from the group consisting of human rhinovirus, herpes simplex I virus, herpes simplex II virus, viral myocarditis and HTLV III virus (AIDS).

19. The method of claim 13 wherein a human is treated for warts.

20. The method of claim 13 wherein a feline species is treated for feline leukemia virus or feline infectious peritonitis.

21. The method of claim 15 wherein a canine species is treated for canine parvovirus.

22. The method of claim 13 wherein canine species is treated for canine herpesvirus.

23. The method of claim 13 wherein a human suffering from HTLV III virus is treated with human alpha-interferon or human beta-interferon.

24. The method of claim 23 wherein human alpha-interferon is administered in aqueous solution at a dosage of about 0.5 to about 1.5 IU/lb per day and wherein said solution of interferon is held in the patient's mouth for a period of time sufficient to allow contact of the interferon by the oral mucosa of said patient.

25. The method of claim 18 wherein the dosage of interferon is divided and administered as a multiple dose daily regimen.

26. The method of claim 23 wherein beta-interferon is administered at a dosage of about 0.5 to about 1.5 IU/lb per day.

27. The method of claim 4 wherein human alpha-interferon is administered at a dosage of about .1 to about 1.5 IU/lb of body weight per day to treat asthma.

28. Method of improving human skin complexion consisting essentially of the administration of interferon at a dosage of about 0.01 to about 5 IU/lb per day in a dosage form adapted to promote contact of said dosage of interferon with the oral and pharyngeal mucosa of a patient suffering a condition of poor skin complexion.

29. An immunotherapeutic oral dosage form of interferon for human use, said dosage form being formulated to provide about 0.01 to about 5 IU/lb (0.022 to about 2.2 IU/kg) of interferon, upon per os administration, for contact with oral and pharyngeal mucosa to stimulate an immunotherapeutic response.

30. The dosage form of claim 29 in the form of a mouthwash.

31. The interferon dosage form of claim 29 in the form of a lozenge adapted to be dissolved during contact with saliva in the mouth.

32. The lozenge of claim 31 formed as a compressed tablet containing about 1 to about 1500 IU of human alpha-interferon or human beta-interferon.

33. A chewable tablet in accordance with claim 31.

34. The interferon dosage form of claim 29 in the form of a syrup containing about 1 to about 1500 IU of human interferon per tablespoon.

35. The immunotherapeutic dosage form of claim 29 wherein the interferon is human alpha-interferon and said dosage form is formulated to provide from about 10 to about 1500 units of alpha-interferon per dose.

36. A method for manufacturing a unitary dosage composition of interferon for use in a method for immunomodulation in a warm-blooded vertebrate, said method of manufacture characterized by combining interferon and a pharmaceutically acceptable carrier therefor adapted to promote contact of about 1 to about 1500 IU of interferon with the oral and pharyngeal mucosa of said vertebrate.

37. The method of claim 36 wherein the interferon is alpha-interferon or beta-interferon.

38. The method of claim 37 wherein the composition is in the form of a lozenge adapted to be dissolved in the mouth upon contact with saliva.

39. An immune system modulating composition containing interferon and a pharmaceutically acceptable carrier therefor, said composition formulated to promote contact of about 1 to about 1500 IU of interferon with the oral and pharyngeal mucosa of an animal or patient in need of immune system modulation.

40. The composition of claim 39 wherein the interferon is alpha-interferon or beta-interferon.

41. The composition of claim 40 in the form of a syrup.

42. The composition of claim 40 in the form of a lozenge adapted to be dissolved in the mouth upon contact with saliva.

43. The composition of claim 39 wherein the interferon is of bovine origin.

AMENDED CLAIMS

[received by the International Bureau on 04 May 1988 (04.05.88);
original claims 1-43 replaced by amended claims 1-44; new claims 45-50 added (7 pages)]

1. In a method utilizing interferon for treatment of an auto immune disorder characterized by a chronic tissue degenerative inflammatory condition in a warm-blooded vertebrate, the improvement which consists essentially of contacting interferon at a dosage of about 0.01 to about 5 IU/lb (about 0.022 to about 11 IU/kg) of body weight per day with the oral and pharyngeal mucosa of said vertebrate.
2. The improvement of claim 1 wherein the administered interferon is human alpha-interferon or human beta-interferon.
3. The improvement of claim 2 wherein the interferon is administered at a dosage of about 0.1 to about 4.0 IU/lb (about 0.22 to about 8.8 IU/kg) per day.
4. The improvement of claim 1 wherein the autoimmune disorder is selected from the group consisting of multiple sclerosis, rheumatoid arthritis, nasal solar dermatitis, stomatitis and lupus erythematosus.
5. The improvement of claim 4 wherein human alpha-interferon is administered to a human patient at a dosage of about 0.5 to about 1.5 IU/lb (about 1.1 to about 3.3 IU/kg) daily until the patient becomes asymptomatic of the autoimmune disorder.
6. The improvement of claim 1 wherein the administered interferon is human beta-interferon.

7. The improvement of claim 4 wherein human interferon is administered to a human patient as an aqueous solution in a dosage of about 0.5 to about 1.5 IU/lb (about 1.1 to about 3.3 IU/kg) per day and wherein said solution of interferon is held in the patient's mouth for a period of time sufficient to allow contact of the interferon by the oral mucosa of said patient.

8. In a method utilizing interferon for treatment of neoplastic disease in a warm-blooded vertebrate, the improvement consisting essentially of contacting said interferon at a dosage of about 0.01 to about 5 IU/lb (about 0.022 to about 11 IU/kg) of body weight per day with the oral and pharyngeal mucosa of said vertebrate.

9. The improvement of claim 8 wherein neoplastic disease in humans is treated by administering human alpha-interferon or human beta-interferon.

10. The improvement of claim 7 wherein alpha-interferon is administered at a dosage of about 0.5 to about 1.5 IU/lb (about 1.1 IU to about 3.3 IU/kg) per day.

11. The improvement of claim 8 wherein the neoplastic disease is selected from the group consisting of malignant lymphoma, melanoma, mesothelioma, Burkitt lymphoma and nasopharyngeal carcinoma.

12. The improvement of claim 8 wherein the neoplastic disease is selected from the group consisting of Hodgkin's disease and leukemia.

13. The improvement of claim 8 wherein the dosage of interferon is divided and administered in a multiple-dose daily regimen.

14. The improvement of claim 8 wherein neoplastic disease in humans is treated by administering human beta-interferon.

15. An improved method of treatment of infectious disease of viral origin in human, canine and feline species consisting essentially of contacting interferon at a dosage of about 0.01 to about 5 IU/lb (about 0.22 to about 11 IU/kg) of body weight per day with the oral and pharyngeal mucosa of said species.

16. The method of claim 15 wherein the dosage is about 0.1 to 4.0 IU/lb (about 0.22 to about 8.8 IU/kg) per day.

17. The method of claim 15 wherein the interferon is selected from alpha-interferon and beta-interferon.

18. The method of claim 15 wherein the interferon is human alpha-interferon.

19. The method of claim 15 wherein human alpha-interferon is administered at dosage of about 0.5 to 1.5 IU/lb (about 1.1 to about 3.3 IU/kg) of body weight per day.

20. The method of claim 19 wherein a human is treated for a viral infection selected from the group consisting of human rhinovirus, herpes simplex I virus, herpes simplex II virus, viral myocarditis and HTLV III virus (AIDS).

21. The method of claim 15 wherein a human is treated for warts.

22. The method of claim 15 wherein a feline species is treated for feline leukemia virus or feline infectious peritonitis.

23. The method of claim 17 wherein a canine species is treated for canine parvovirus.

24. The method of claim 15 wherein canine species is treated for canine herpesvirus.

25. The method of claim 15 wherein a human suffering from Acquired Immune Deficiency Syndrome is treated with human alpha-interferon or human beta-interferon.

26. The method of claim 25 wherein human alpha-interferon is administered in aqueous solution at a dosage of about 0.5 to about 1.5 IU/lb (about 1.1 to about 3.3 IU/kg) per day and wherein said solution of interferon is held in the patient's mouth for a period of time sufficient to allow contact of the interferon by the oral mucosa of said patient.

27. The method of claim 20 wherein the dosage of interferon is divided and administered as a multiple dose daily regimen.

28. The method of claim 25 wherein beta-interferon is administered at a dosage of about 0.5 to about 1.5 IU/lb (about 1.1 to about 3.3 IU/kg) per day.

29. A method of treating a condition characterized by a hyperallergenic response to antigens, said method comprising the administration of interferon at a dosage of about 0.01 to about 5 IU/lb (0.022 to about 11 IU/kg) per day in a dosage form adapted to promote contact of said dosage of interferon with the oral and pharyngeal mucosa of a warm-blooded vertebrate suffering such condition.

30. The method of claim 25 wherein human alpha-interferon is administered at a dosage of about 0.1 to about 1.5 IU/lb (about 0.22 to about 3.3 IU/kg) of body weight per day to treat asthma.

31. Method of improving human skin complexion comprising the administration of interferon at a dosage of about 0.01 to about 5 IU/lb (about 0.022 to about 11 IU/kg) per day in a dosage form adapted to promote contact of said dosage of interferon with the oral and pharyngeal mucosa of a patient suffering a condition of poor skin complexion.

32. Method of claim 31 wherein human alpha-interferon is administered to treat acne.

33. Method of treating bacterial infections in warm-blooded vertebrates, said method comprising administering to said vertebrate interferon at a dosage of about 0.01 to about 5 IU/lb (about 0.022 to about 11 IU/kg) per day in a dosage form adapted to promote contact of said dosage of interferon with the oral and pharyngeal mucosa of said vertebrate.

34. The method of claim 33 wherein human alpha-interferon is utilized to treat warm-blooded vertebrates infected with antibiotic resistant microorganisms.

35. The method of claim 34 wherein the warm-blooded vertebrate is a human.

36. The method of claim 35 wherein human alpha-interferon is administered at a dosage of about 0.5 to about 1.5 IU/lb (about 1.1 to about 3.3 IU/kg) per day.

37. An improved immunotherapeutic oral dosage form of interferon for human use, said dosage form being formulated so that each unit dose of said dosage form provides about 0.01 to about 5 IU of interferon/lb (about 0.022 to about 11 IU/kg) of body weight for contact with oral and pharyngeal mucosa to stimulate an immuno therapeutic response.

38. The dosage form of claim 37 in the form of a mouthwash.

39. The interferon dosage form of claim 37 in the form of a lozenge adapted to be dissolved during contact with saliva in the mouth.

40. The lozenge of claim 39 formed as a compressed tablet containing about 1 to about 1500 IU of human alpha-interferon or human beta-interferon.

41. A chewable tablet in accordance with claim 37.

42. An effervescent tablet in accordance with claim 37 adapted for dissolution in water to form a mouthwash or gargle formulation.

43. The interferon dosage form of claim 37 in the form of a syrup containing about 1 to about 1500 IU of human interferon per tablespoon.

44. The immunotherapeutic dosage form of claim 37 wherein the interferon is human alpha-interferon and said dosage form is formulated to provide from about 10 to about 1500 IU of alpha-interferon per dose.

45. A method for manufacturing a composition for use in an improved method of using interferon to stimulate an immunotherapeutic response in a

warm-blooded vertebrate, said method of manufacture characterized by combining interferon and a pharmaceutically acceptable carrier therefor to form an interferon composition adapted to provide, upon unit dose administration, about 0.01 to about 5 IU of interferon per lb (about 0.022 to about 11 IU/kg) of body weight for contact essentially with the oral and pharyngeal mucosa of said vertebrate to stimulate an immunotherapeutic response.

46. The method of claim 45 wherein the interferon is alpha-interferon or beta-interferon.

47. The method of claim 45 wherein the interferon is alpha-interferon.

48. The method of claim 45 wherein the pharmaceutically acceptable carrier is selected so that the composition can be held in the mouth of the warm-blooded vertebrate for a time sufficient to allow contact of the interferon with the oral and pharyngeal mucosa of said vertebrate.

49. The method of claim 45 wherein the pharmaceutically acceptable carrier is selected so that the composition is in a solid dosage form.

50. The method of claim 45 wherein the pharmaceutically acceptable carrier is selected so that the composition is in a liquid dosage form.

INTERNATIONAL SEARCH REPORT

International Application No. PCT/US87/02998

I. CLASSIFICATION OF SUBJECT MATTER (If several classification symbols apply, indicate all) ⁶

According to International Patent Classification (IPC) or to both National Classification and IPC

IPC (4) : A61K 45/02

U.S.CI. : 424/85

II. FIELDS SEARCHED

Minimum Documentation Searched ⁷

Classification System	Classification Symbols
U.S.	424/85

Documentation Searched other than Minimum Documentation
to the Extent that such Documents are Included in the Fields Searched ⁸

Online Computer Search of Chemical Abstracts 1967-1988
Search terms: interferon, oral administration

III. DOCUMENTS CONSIDERED TO BE RELEVANT ⁹

Category ¹⁰	Citation of Document, ¹¹ with indication, where appropriate, of the relevant passages ¹²	Relevant to Claim No. ¹³
Y	US, A, 4,497,795 (CUMMINS) 05 February 1985. See entire document.	1-43
Y	US, A, 4,462,985 (CUMMINS) 31 July 1984. See entire document.	1-43
Y	The Lancet, Volume 1, issued May 1979 (U.S.A.), Kajander, "Interferon Treatment of Rheumatoid Arthritis". See pages 984-985.	1-43
Y	J. Interferon Res., Volume 6, issued June 1986 (U.S.A.), Yamamoto, "Human Alpha and Beta Interferon But Not Gamma-Suppress the In Vitro Replication of LAV, HTLV-III, and ARV-2". See page 143-152.	1-43
Y	FR, A, 2575655 (CUMMINS) 28 July 1986. See entire document.	1-43

- Special categories of cited documents: ¹⁰
- "A" document defining the general state of the art which is not considered to be of particular relevance
- "E" earlier document but published on or after the International filing date
- "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified)
- "O" document referring to an oral disclosure, use, exhibition or other means
- "P" document published prior to the International filing date but later than the priority date claimed

- "T" later document published after the International filing date or priority date and not in conflict with the application but cited to understand the principle or theory underlying the invention
- "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to involve an inventive step
- "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such documents, such combination being obvious to a person skilled in the art.
- "&" document member of the same patent family

IV. CERTIFICATION

Date of the Actual Completion of the International Search

14 MARCH 1988

International Searching Authority

ISA/US

Date of Mailing of this International Search Report

12 APR 1988

Signature of Authorized Officer

Blondel Hazel

Blondel Hazel